Appl. No. 10/717,236 Amendment dated February 25, 2010 Reply to Office action dated February 19, 2010

This listing of claims will replace all prior versions and listings of claims in the application:

Listing of Claims

 (Previously Presented) A method for preparing lithium transitional metal oxides, comprising the steps of:

preparing a carbonate precursor using the following substeps:

carbonate precursor, Ni_{1-x-v}Co_xMn_vCO₃; and

forming a first aqueous solution containing a mixture of at least two of the ions of the following metal elements ("Meⁿ"): cobalt (Co), nickel (Ni), and manganese (Mn); forming a second aqueous solution containing ions of CO₃²; and mixing and reacting said first solution and said second solution to produce the

preparing said lithium transition metals oxide from said carbonate precursors using the following substeps:

evenly mixing ${\rm Li}_2{\rm CO}_3$ and said carbonate precursor; calcinating the mixed material in high temperature; and cooling and pulverizing the calcinated material to obtain said lithium transition metal oxide, ${\rm Li}\ Ni_{1\times N}{\rm Co}_3{\rm Mn},{\rm O}_2$.

 (Previously Presented) The method of claim 1 wherein the calcinating step is performed in air. Appl. No. 10/717,236 Amendment dated February 25, 2010 Reply to Office action dated February 19, 2010

3. (Previously Presented) The method of claim 1 further comprising a step after the cooling

step, again calcinating said cooled and pulverized material in air at high temperature.

4. (Previously Presented) The method of claim 3 further comprising a step after the again

calcinating step, cooling, ballmilling, and sifting the again calcinated material to obtain the

compound of lithium transition metal oxide, Li Ni1-x-yCoxMnyO2.

5. (Previously Presented) The method of claim 1 wherein the mixing and reacting substep

is performed under the condition where the PH is between 5 to 10.

6. (Previously Presented) The method of claim 1 wherein the mixing and reacting substep

is performed under the condition where the reaction temperature is between 30 degrees to 90

degrees Celsius.

(Previously Presented) The method of claim 1 wherein the mixing and reacting substep

is performed under the condition where the stirring speed is between 20 to 120 revolutions per

minute.

8. (Previously Presented) The method of claim 1 wherein the mixing Li₂CO₃ substep is

performed under the condition where the calcination is at 500 to 800°C.

Appl. No. 10/717,236 Amendment dated February 25, 2010

Amendment dated February 25, 2010 Reply to Office action dated February 19, 2010

9. (Previously Presented) The method of claim 3 wherein the again calcinating step is

performed under the condition where the calcination is at 700 to 950°C.

10. (Previously Presented) The method of Claim 1 wherein the ion concentration of the first

solution is 0.1 to 3.0 mol/L

11. (Previously Presented) The method of claim 10 wherein the Meⁿ⁺ derives from at least

one of the compounds from the following groups of materials: sulfates, nitrates, chlorides.

acetates, other water soluble inorganic salts, and, other water soluble salts of organic acid.

12. (Previously Presented) The method of claim 1 wherein the first solution contains at least

one type of additive ingredient from the group containing the ions of the following elements: Ca,

Mg, Zn, Y, Ga, and In.

13. (Previously Presented) The method of claim 12 wherein the molar concentration of the

additive ingredient is of 0 to 10% of the overall solution.

14. (Previously Presented) The method of claim 1 wherein the CO₃²⁻ ion concentration of the

second solution is 0.1 to 3.0 mol/l.

15. (Previously Presented) The method of claim 1 further including a third solution in the

preparation of the carbonate precursor wherein the third solution is a NaOH solution; and

wherein the condition for preparing said carbonate precursor is where the flow rate is adjusted

for the reaction to proceed at PH between 8 to 9 and the temperature is between 30-90°C.

16. (Previously Presented) The method of claim 1 wherein said first solution is an aqueous

solution of the combination of the sulfates salts of cobalt, nickel, and manganese; and said

second solution is (NH₄)₂CO₃; and further including a third solution that is a NH₂·H₂O solution

in the preparation of said carbonate precursor where the flow rate is adjusted for the reaction to

proceed at the PH level between 8 to 9 and the temperature is between 30 to 90°C.

17. (Previously Presented) The method of claim 1 wherein the mixing the Li₂CO₂ step is to

first wet mixing with an organic solvent such as ethanol.

18. (Previously Presented) A method for preparing lithium transitional metal oxides,

comprising the steps of:

preparing a carbonate precursor using the following substeps:

forming a first aqueous solution containing a mixture of at least two of the ions of

the following metal elements ("Meⁿ⁺"): cobalt (Co), nickel (Ni), and manganese (Mn),

wherein the ion concentration of the first solution is 0.1 to 3.0 mol/l; wherein the first

solution contains at least one type of additive ingredient from the group containing the

ions of the following elements: Ca, Mg, Zn, Y, Ga, and In; and wherein the molar

concentration of the additive ingredient is of 0 to 10% of the overall solution;

forming a second aqueous solution containing ions of CO₃²; and

mixing and reacting said first solution and said second solution to produce the

carbonate precursor, Ni_{1-x-v}Co_xMn_vCO₃, under the conditions where the PH is between 5

to 10, the reaction temperature is between 30 degrees to 90 degrees Celsius, and the

stirring speed is between 20 to 120 revolutions per minute; and

preparing said lithium transition metals oxide from said carbonate precursors using the

following substeps:

evenly mixing Li₂CO₃ and said carbonate precursor;

calcinating the mixed material in air in high temperature of 500 to 800°C; and

cooling and pulverizing the calcinated material;

again calcinating said cooled and pulverized material in air at high temperature of

700 to 950°C; and

cooling, ballmilling, and sifting the again calcinated material to obtain the

compound of lithium transition metal oxide, Li $Ni_{1\text{-x-y}}Co_xMn_yO_2$.

19. (Previously Presented) The method of claim 18 wherein the Meⁿ⁺ derives from at least

one of the compounds from the following groups of materials: sulfates, nitrates, chlorides,

acetates, other water soluble inorganic salts, and, other water soluble salts of organic acid.

20. (Previously Presented) The method of claim 18 wherein the CO₃²⁻ ion concentration of

the second solution is 0.1 to 3.0 mol/L